Eommunications

[16] We found that BRP-H is also a polymerization catalyst for other thiophenes, for example, 3-dodecyl-thiophene.

[17] Handbook of Oligo- and Polythiophenes (Ed.: D. Fichou), Wiley-VCH, Woinheim, 1999.

[18] D. Dini, F. Decker, F. Andreani, E. Salatelli, P. Hapiot, Polymer 2000, 41, 6473.

[19] In the measurement setup an argon ion laser beam was focused by a 25 \times microscope objective to a 50 μm diameter spot on the sample. The fluorescence emission was imaged by a .50 x objective on a CCD camera.

[20] No UV/Vis spectrum could be recorded from a single vesicle, nor from a suspension of vesicles, because of scattering problems.

[21] A 0.5 gL-1 solution of PS-PIAT dissolved in THF was injected into a 30 mg L-1 solution of CAL B enzymes; the final water/ THF ratio was 12:1 (v/v). After the mixture was left for two days to equilibrate, it was dialyzed to dispose of all nonincluded cazymes.

Somatostatin Mimics

Design and Synthesis of γ-Dipeptide Derivatives with Submicromolar Affinities for Human Somatostatin Receptors

Dieter Seebach,* Laurent Schaeffer, Meinrad Brenner, and Daniel Hoyer

In a previous paper we have shown that simple N-acyl-ydipeptide amides that resemble a βII' turn of an α-peptide can be designed to form a turn structure in solution (NMR) and in the solid state (X-ray).[1,2] To see whether such a turn could also be used to mimic a peptide, the biological activity of which rests upon a turn structure carrying functionalized side chains, we have now synthesized compounds 12-g (Scheme 1), with the side chain of tryptophan in the y2 position of the first and of lysine in the Y position of the second y-amino acid, and have tested their affinities for the human somatostatin receptors hsst₁₋₅.[3-6]

The synthesis of y-dipeptide derivatives 1 commenced with the N-Boc-y-lactams 2 and 3 (Boc = tert-butoxycarbonyl), readily available from the corresponding commercial (R)-Ala and (S)-Lys acids by known procedures. [1,7] Ring opening (with the Lys derivative after change of side-chain protection, -4), and esterification with Me3Si(CH2)2OH provided the (R)-Boc-Y-hhAla and Boc-Y-hhLys(Bn2) esters, which were

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Dr. D. Høyer Novartis Pharma AG Nervous System Research S-386-745, 400Z Basel (Switzerland)



NR^ZR³

RZ H_1 . NR²E³ Βn Βn Bα Bn Bn Me н Mie B٥ Ðη н Me Nap н н н Mes Expected conformation of 1

Scheme 1. Structural formulae of the y-peptides 1. In the expected conformation of 1 the red arrow points to a CH2 group of the H2N(CH3)4 unit, which is placed inside the shielding cone of the aromatic incole ring. Mes = mesitylenesulfonyl, Bn = benzyl, Nap = naphthyl.

5, $H = (CH_z)_z SIMe_2$ 6, H = H

NBn₂
7.
$$R^1 = Boc$$
, $R^2 = Mc$, $X = O(CH_2)_2SiMe_3$
8. $R^1 = R^2 = H$, $X = NHMc$
9. $R^1 = H$. $R^2 = Mc$, $X = NHMc$

doubly deprotonated and alkylated with 1-mesitylenesullonyl-3-bromomethylindole and Mel to give the unlike 71-amino acid derivatives 5 and 7, respectively. The ester group in compound 5 with Trp side chain was cleaved (Bu,NP, -6). and the lysine-derived esters were converted to the methylamides 8 and 9 without and with 2-methyl substitution, respectively (1. Bu, NF, 2. MeNH₂, 3. F₃CCO₂H). Coupling of the two y-amino acid derivatives (6 + 8 and 6 + 9), removal of the Boc groups, and acylation with 2-naphthylacetic acid¹⁵ (4-methylmorpholine, 1-hydroxy-1H-benzotriazole, 1-cthyl: 3-(3-dimethylaminopropyl)carbodiimide) produced the sidechain-protected N-acyl-dipeptide amides 1a and 1b. Deprotection procedures (MeSO₃H, F₃CCO₂H, and Pd/C, H₂) led to the various partially or fully deprotected y-dipeptide derivatives 1c-1g. All compounds were purified and fully characterized by elemental analyses, specific optical rotations circular dichroism (CD), IR, and NMR spectroscopy. 20d mass spectrometry.

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Figure 1. H peptides 14 **feld-shifted**

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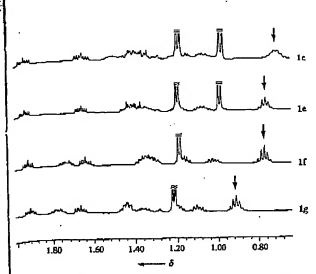


figure 1. High-field part of the 500 MHz 1H NMR spectra of the Y-dipeptides 1c, 1c, 1f, and 1g in CD,OD. The red arrows point to highfeld-shifted N(CH2)4 resonances.

A typical feature of the turn structure in somatostatin and its analogues is the juxtaposition of the tryptophan and lysine ade chains, which places CH2 groups of the H2N(CH2)4 unit inside the shielding cone of the aromatic indole ring (NMR hifts between $\delta = 0.8$ and 0.3 ppm are observed). [9] High-field sections of the NMR spectra of four y-dipeptide amides,

shown in Figure 1, in which CH2 signals appear between $\delta = 0.9$ and 0.6 ppm, confirm the proximity beween the corresponding side chains and are thus compatible with a turn conformation of these monounds. The CD spectra of the N-naphthylacetyl dipeptide amides lexhibit an intensive negative Cotion effect near 200 nm ([Θ] up to $70000 \text{ deg cm}^2 \text{dmol}^{-1}$), with

weaker and broader peak near 220 nm ([$oldsymbol{ heta}$] up to 30000 deg cm2 dmol-1) (Figure 2); this CD pattern may be taken as another piece of evidence for the presence of a sccondary structure.

Probably the most stringent test of the y-dipeptide structure is the affinity for somatostatin receptors. Binding affinities for the five cloned human receptors hsst₁₋₅, expressed in CCL-39 cell lines, were determined by displacement of [125]]LTT-SRIF28 from these receptor proteins [10] While the fully protected γ -dipeptide 1d binds to hsst₁ and hsst, with remarkable K_D values of 0.55 and 1.00 μM, respectively, the partially and the fully deprotected y-dipeptide derivatives 1 f and 1g bind to hsst₅ with K_0 values of 0.51 and 0.87 μμ, respectively (Table 1). Intriguingly, the highest affinities (1d/hsst, 11/hsst) are observed when the side chain functional groups (3-indolylmethyl and (CH2)4NH3+) are protected by bulky aromatic moicties (N-mesitylenesulfonyl and/or -benzyl)!

The results presented here are confirmative, surprising, and promising; they demonstrate that a 14-amino-acid cyclic disulfide hormone, somatostatin, can be mimicked by a simple, designed, low-molecular-weight, open-chain y-dipeptide derivative (cf. 1g) that contains only three amide bonds; they suggest that hitherto unknown hydrophobic pockets are present in the receptors (hsst₁, hsst₂, and hsst₅), which supposedly house the turn-bound Trp and Lys side chains (cf. 1c, 1d, 1f); and they promise a potential of γ -peptides for the development of peptidase-resistanting peptidomimetic

Table 1: ρK_D Values for γ-peptides 1 b-1 g at the five hast receptors expressed in CCL-39 cells and wedioligand hinding assays with [125][LTT-SRIF as radioligand. PLOT

Receptor	16	1c	14	le	1 f	٦g `	Octreotide ^(b)	5RIF ₁₄ I
hsst _i	5.47	6.06	6.26	5.61	5.98	4.73	6.45	9.08
hsst,	<5	<5	5.17	<5	5.01	2.81	9.11	10.06
hsst,	5.53	5.89	6.00	5.73	5.67	5.42	8.60	9.67
hsst,	4.67	5.74	5.92	5.66	5.79	5.44	5.76	8.39
hsst.	4.49	5.01	5.87	5.14	6.29	6.06	7.31	9.01

[a] Submicromolar affinities are highlighted in red. [b] Sandostatin. (c) Somatostatin.

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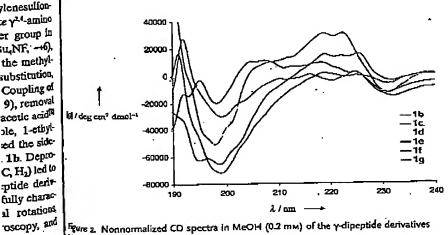


Figure 2. Nonnormalized CD spectra in MeOH (0.2 mm) of the y-dipeptide derivatives

Received: September 25, 2002 [Z50242]

[1] M. Brenner, D. Seebach, Helv. Chim. Acta **2001, 84, 2155–2166**.

[2] For a non-peptidic analogue, see: R. W. Hoffmann, Angew. Chem. 2000, 112, 2134-2150; Angew. Chem. Inc. Ed. 2000, 39, 2054-2070.

[3] For \(\beta\)-tetrapoptides mimicking, somatostatin with nanomolar affibities for hsst, see: K. Gademann, M. Ernst, D. Hoyer, D. Seebach, Angew. Chem. 1999, 111, 1302-1304; Angew. Chem. Int Ed. 1999, 38, 1223-1226; K. Gademann, M. Ernst, D. Seebach, D. Hoyer, Helv. Chim. Acta 2000, 83, 16-33; K. Gademann, T. Kimmerlin, D. Hoyer, D. Seebach, J. Med. Chem. 2001, 44, 2460-2468; D. Scebach, M. Rueping, P. J. Arvidsson, T. Kimmerlin, P.

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Communications

Micuch, C. Noti, D. Langenegger, D. Hoyer, Helv. Chim. Acta 2001, 84, 3503-3510.

- [4] For a review on somatostatin receptors, see: D. Hoyer, H. Luebbert, C. Bruns, Naunyn-Schmiedeberg's Arch. Pharmacol. 1994, 350, 441-453; J. P. Hannon, C. Nunn, B. Stolz, C. Bruns, G. Weckbecker, I. Lewis, T. Troxler, K. Hurth, D. Hoyer, J. Mol. Neurosci. 2002, 18, 15-27. Classification and nomenclature of somatostatin receptors: D. Hoyer, G. I. Bell, M. Berelowitz, J. Epelbaum, W. Feniuk, P. P. Humphrey, A. M. O'Carroll, Y. C. Patel, A. Schonnbrunn, J. E. Taylor, T. Reisine, Trends Pharmacol. Sci. 1995, 16, 86-88.
- [5] For a review on somatostatin analogues, see: A. Janecka, M. Zubrzycka, T. Janecki, J. Pept. Res. 2001, 58, 91-107.
- [6] Methyl groups in the γ' position of the first and in the γ' position of the second γ-amino acid do stabilize the turn (see ref. [1] for more details).
- M. Smreina, P. Majer, E. Majerova, T. A. Guerassina, M. A. Eissanstat, Tetrahedron 1997, 53, 12867-12874; S. Hanessian, R. Schaum, Tetrahedron Lett. 1997, 38, 163-166; T. Hintermann, K. Gademann, B. Jaun, D. Seebach, Helv. Chim. Acta 1998, 81, 983-1002.
- [8] The choice of a 2-naphthylacetyl group at the N-terminus was inspired by previous observations: The naphthyl group may be considered to mimic the benzyl side chain of a Phe residue, which precedes the Trp in the peptide chain of somatostatin: S. J. Hocart, R. Jain, W. A. Murphy, J. E. Taylor, B. Morgan, D. H. Coy, J. Med. Chem. 1998, 41, 1146-1154; A. J. Souers, A. A. Virgilio, A. Rosenquist, W. Fenuik, J. A. Ellman, J. Am. Chem. Soc. 1999, 121, 1817-1825.
- [9] B. H. Arison, R. Hirschmann, D. F. Veber, Bioorg. Chem. 1978, 7, 447-451; R. M. Freidinger, D. S. Perlow, W. C. Randall, R. Saperstein, B. H. Arison, D. F. Veber, Int. J. Pept. Protein Res. 1984, 23, 142-150; C. Wynants, D. Tourwe, W. Kazmierski, V. I. Hruby, G. Van Binst, Eur. J. Biochem. 1989, 185, 371.
- [10] J. P. Hannon, C. Petrucci, D. Fehlmann, C. Viollet, J. Epelbaum, D. Hoyer, Neuropharmacology 2002, 42, 396-413. CCL-39 stands for Chinese hamster lung fibroblast cells No. 39. [125] LTT-SRIF₂₈ is a somatostatin dimer containing three mutations (Leu⁸, p-Trp², ¹²⁵I-Tyr), and SRIF is the acronym for somatostatin release inhibiting factor, 28 refers to the numbere of amino acids in the physiologically active form of the hormone.
- [11] J. Franckenpohl, P. I. Arvidsson, J. V. Schreiber, D. Seebach, ChemBioChem 2001, 2, 445-455.

Controlled Electropolymerization:



Liquid-Crystal Templating of Conducting Polymers**

James F. Hulvat and Samuel I. Stupp*

In organic electronics, conducting polymers have a number of advantages over small molecules, particularly because of their stability, mechanical properties, and case of processing. However, performance of some conducting polymers is limited by their high degree of disorder. And Molecular ordering improves carrier mobility in organic field-effect transitors and enhances charge injection in organic light-emitting diodes (OLEDs). For this reason, vapor-sublimated crystalline films of small molecules are often used, but alternative strategies to obtain molecular ordering would reduce cost and simplify fabrication of organic electronic devices. One possible way to achieve this is through molecular self-organization. Toward this goal we developed an aqueous low-temperature technique for preparing conducting polymer films in a self-organized template.

Films of poly(3,4-ethyldioxythiophene) (PEDOT) are commonly used as hole injection layers in OLEDs PEDOT can be polymerized in organic solvents or in an aqueous suspension with a soluble copolymer or surfactant, leading to amorphous films. We have studied here the formation of PEDOT films by electropolymerization within a liquid crystalline template.

The well-known hexagonal (H1) lyotropic liquid crystal (LC) consists of cylindrical hydrophobic cores parallel to one another and separated by a hydropholic continuum (see Figure 1 in the Supporting Information). LCs have been used by us and others to template inorganic minerals as well as in the formation of mesoporous silica. With regard to conducting polymers, template chemistry has been used to incorporate chains within the channels of mesoporous silica. These approaches, however, are limited to soluble, chemically polymerized polymers or to water-soluble monomers such as aniline or pyrrole. Our approach described here is novel in two key respects. First, polymerization occurs in the bydrophobic domain of the LC, allowing use of less polar

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^[***] This work made use of the Electron Probe Instrumentation Center at Northwestern University and was funded by a DoE grant (DE-FG02-Northwestern University and was funded by a NOSEG fellowship. We ODER45810/A001). J.F.H. Is supported by an NOSEG fellowship. We thank M. Kern for assistance with XRD and Prof. I. Koltover for balance discussions.

helpful discussions.

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